# Phylogeny and taxonomy of the genus Cylindrocladiella 

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#### Abstract

The genus Cylindrocladiella was established to accommodate Cylindrocladium-like fungi that have small, cylindrical conidia and aseptate stipe extensions. Contemporary taxonomic studies of these fungi have relied on morphology and to a lesser extent on DNA sequence comparisons of the internal transcribed spacer regions (ITS 1, 2 and 5.8S gene) of the ribosomal RNA and the $\beta$-tubulin gene regions. In the present study, the identity of several Cylindrocladiella isolates collected over two decades was determined using morphology and phylogenetic inference. A phylogeny constructed for these isolates employing the $\beta$-tubulin, histone H3, ITS, 28S large subunit and translation elongation factor 1-alpha gene regions resulted in the identification of several cryptic species in the genus. In spite of


[^0]the 18 new Cylindrocladiella species described in this study based on morphological and sequence data, several species complexes remain unresolved.

Keywords Cylindrocladiella $\cdot$ Cryptic species • Phylogeny • Taxonomy

## Introduction

The genus Cylindrocladiella was established by Boesewinkel (1982) to accommodate five Cylindrocladium-like species producing small, cylindrical conidia. Cylindrocladiella, which is based on C. parva, is distinguished from the anamorph state of Calonectria (= Cylindrocladium) by its symmetrically branched conidiophores that can be penicillate and/or subverticillate, producing an asymmetrical bundle of small, cylindrical, 1-septate conidia ( $<20 \mu \mathrm{~m}$ in length), aseptate stipe extensions, and having Nectricladiella teleomorphs (Boesewinkel 1982, Crous and Wingfield 1993, Schoch et al. 2000). The Nectricladiella teleomorphs are characterised by their perithecia having smooth walls that collapse laterally when dry, and brown setae arising from the perithecial wall surface (Schoch et al. 2000).

Initially, the generic status of Cylindrocladiella was strongly contested (Peerally 1991, Sharma and Mohanan 1991). Morphological evaluations and comparisons by Crous and Wingfield (1993) and Crous et al. (1994), however, confirmed the generic status of this genus, which was later supported by molecular data (Victor et al. 1998, Schoch et al. 2000). Victor et al. (1998) used RFLPs, ATDNA data and morphological comparisons, to recognise seven species in the genus. This was later supported by phylogenetic inference of the ITS and partial $\beta$-tubulin gene regions, resulting in the addition of another species to the
genus (Schoch et al. 2000). To date, nine species of Cylindrocladiella are recognized, with only two connected to their respective Nectricladiella teleomorph states (Crous 2002, van Coller et al. 2005).

Cylindrocladiella spp. are soil-borne fungi, and are generally regarded as pathogens and/or saprobes of various plant hosts and substrates in temperate, sub-tropical and tropical regions worldwide (Crous 2002, van Coller et al. 2005, Scattolin and Montecchio 2007). They have been associated with a variety of disease symptoms that included leaf spots, and rots of roots, stems and cuttings of agricultural, forestry and horticultural crops (Crous et al. 1991, Peerally 1991, Crous and Wingfield 1993, Victor et al. 1998, Crous 2002, van Coller et al. 2005, Scattolin and Montecchio 2007).

The aim of this study was to consider the identity of several Cylindrocladiella isolates collected over the past two decades from various substrates and regions of the world. To achieve this goal, morphological and culture characteristics were combined with multigene phylogenetic inference for all isolates studied.

## Material and methods

## Isolates

Isolates and ex-type strains of Cylindrocladiella spp. were obtained from various culture collections, isolated from symptomatic plant material and/or baited from soils as described in Crous (2002) and indicated in Table 1. Representative strains have been maintained in the culture collection of the CBSKNAW Fungal Biodiversity Centre (CBS) and the working collection of Pedro Crous (CPC).

## Phylogeny

Total genomic DNA was extracted from 7-10 day old single-conidial cultures using the technique described by Möller et al. (1992). Partial fragments of the following genes and gene regions were amplified using the PCR conditions and primer sets mentioned in Lombard et al. (2010b): $\beta$-tubulin (BTUB), histone H3 (HIS3), internal transcribed spacers and 5.8 s rDNA (ITS), 28 s large subunit (LSU) and translation elongation factor 1-alpha (TEF1- $\alpha$ ). The PCR reactions were carried out using a MyCycler ${ }^{\text {TM }}$ thermal cycler (Bio-Rad Laboratories, Inc.) consisting of an initial step of $95^{\circ} \mathrm{C}$ for 5 min followed by 40 cycles of 30 s at $95^{\circ} \mathrm{C}, 30 \mathrm{~s}$ at $52^{\circ} \mathrm{C}, 1 \mathrm{~min}$ at $72^{\circ} \mathrm{C}$ and ending with a final extension step of 7 min at $72^{\circ} \mathrm{C}$.

Amplicons were sequenced in both directions using the same primer sets used for amplification and the consensus sequences were aligned using MAFFT v6.611 (Katoh and

Toh 2008) for each gene region. Ambiguous regions in the alignments were removed manually and both ends of the sequences were truncated. All sequences obtained were deposited in GenBank with accession numbers listed in Table 1.

Analyses of the DNA sequence data were done using PAUP (Phylogenetic Analysis Using Parsimony, v4.0b10, Swofford 2002). Initial neighbour-joining analyses (NJ) with the uncorrected ("p"), Juke-Cantor and HKY85 substitution models were done using the LSU sequence data to determine if the Cylindrocladiella isolates used in this study formed a monophyletic group. Congruency of the sequence data for each locus were determined using visual inspection of the tree topologies of $70 \%$ reciprocal NJ bootstrap trees (Gueidan et al. 2007) determined as described in Lombard et al. (2010c). Thereafter, the combined DNA sequence dataset was subjected to maximum parsimony (MP) and Bayesian analyses.

For the MP analysis, the phylogenetic relationships were determined using a heuristic search with 1000 random sequence additions with a tree bisection-reconnection algorithm and the branch swapping option set on "best trees" only. All characters were weighted equally and alignment gaps were treated as missing data. Measures calculated for parsimony included tree length (TL), consistency index (CI), retention index (RI) and rescaled consistence index (RC). Branch support was assessed using a 1000 bootstrap replicates.

For Bayesian analysis, a Markov Chain Monte Carlo (MCMC) algorithm was used to generate phylogenetic trees with posterior probabilities using MrBayes v3.1.1 (Ronquist and Heulsenbeck 2003). Nucleotide substitution models were determined for each gene using the Akaike Information Criterion (AIC) in MrModeltest v2.3 (Nylander 2004) and included in the analysis. The DNA sequence data was subjected to two separate analyses of four MCMC chains run from random trees for $1,000,000$ generations with sampling at every 100 generations. Both runs converged on the same likelihood score and tree topology, and therefore the first 1,000 trees were discarded as the burn-in phase. Posterior probabilities were determined from the remaining trees. All alignments and trees generated in this study, have been deposited in TreeBASE (http:/www. treebase.org).

## Taxonomy

Single-conidial cultures of Cylindrocladiella isolates were prepared on synthetic nutrient-poor agar (SNA; Nirenburg 1981) as described in Lombard et al. (2009). In some cases, carnation leaf pieces were added to the SNA to promote sporulation. The gross morphological characteristics were determined with 30 measurements at $\times 1,000$ magnification
Table 1 Isolates of Cylindrocladiella studied

| Species | Isolates | GenBank Accessions |  |  |  |  | Substrate | Country | Collector |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | BTUB | HIS3 | ITS | LSU | TEF-1 $\alpha$ |  |  |  |
| C. australiensis | CBS 129567 $=$ CPC 17507 ${ }^{\text {T }}$ | JN098747 | JN098932 | JN100624 | JN099222 | JN099060 | soil | Australia | P.W. Crous |
|  | CBS 129568=CPC 17562 | JN098748 | JN098931 | JN100623 | JN099221 | JN099059 | soil | Australia | P.W. Crous |
| C. camelliae | CPC234=PPRI 3990=IMI 346845 | AY793471 | AY793509 | AF220952 | JN099249 | JN099087 | Eucalyptus grandis | South Africa | P.W. Crous |
|  | CPC 237 | JN098749 | JN098839 | JN100573 | JN099252 | JN099090 | Eucalyptus grandis | South Africa | P.W. Crous |
|  | CPC 239 | JN098750 | JN098838 | JN100571 | JN099250 | JN099088 | Eucalyptus grandis | South Africa | P.W. Crous |
|  | CBS 114891= ${ }^{\text {CPC } 277}$ | AY793472 | AY793510 | AF220953 | JN099248 | JN099086 | Eucalyptus grandis | South Africa | P.W. Crous |
| C. clavata | CBS 129563=CPC 17591 | JN098751 | JN098859 | JN099096 | JN099136 | JN098975 | soil | Australia | P.W. Crous |
|  | CBS 129564 $=$ CPC 17592 ${ }^{\text {T }}$ | JN098752 | JN098858 | JN099095 | JN099135 | JN098974 | soil | Australia | P.W. Crous |
| C. cymbiformis | CBS 129553 $=$ CPC 17393 ${ }^{\text {T }}$ | JN098753 | JN098866 | JN099103 | JN099143 | JN098988 | soil | Australia | P.W. Crous |
|  | CBS 129554=CPC 17392 | JN098754 | JN098867 | JN099104 | JN099144 | JN098989 | soil | Australia | P.W. Crous |
| C. elegans | CBS 338.92=PPRI 4050 $=$ IMI $346847^{\text {T }}$ | AY793474 | AY793512 | AY793444 | JN099201 | JN099039 | leaf litter | South Africa | I. Rong |
|  | CBS 110801=CPC 525 | JN098755 | JN098916 | JN100609 | JN099206 | JN099044 | leaf litter | South Africa | P.W. Crous |
| C. ellipsoidea | CBS 129572=CPC 17558 | JN098756 | JN098943 | JN100636 | JN099235 | JN099073 | soil | Australia | P.W. Crous |
|  | CBS 129573= CPC 17560 ${ }^{\text {T }}$ | JN098757 | JN098857 | JN099094 | JN099134 | JN098973 | soil | Australia | P.W. Crous |
|  | CPC 17559 | JN098758 | JN098942 | JN100635 | JN099234 | JN099072 | soil | Australia | P.W. Crous |
|  | CPC 17561 | JN098759 | JN098853 | JN099093 | JN099133 | JN098972 | soil | Australia | P.W. Crous |
| C. hawaiiensis | CBS 118704 | JN098760 | JN098878 | JN099115 | JN099158 | JN098996 | soil | Hawaii | Y. Degawa |
|  | CBS 129569 $=$ CPC 12272 ${ }^{\text {T }}$ | JN098761 | JN098929 | JN100621 | JN099219 | JN099057 | soil | Hawaii | Y. Degawa |
| C. infestans | CBS 111795=ATCC 44816=CPC $2380^{\text {T }}$ | AF320190 | AY793513 | AF220955 | JN099199 | JN099037 | Pinus pinea | New Zealand | H.J. Boesewinkel |
|  | CBS 191.50=IMI 299376=CPC 2480 | AY793475 | AY793514 | AF220956 | JN099198 | JN099036 | Arenga pinnata | Indonesia | K.B. Boedijn \& J. Reitsma |
|  | CBS 192.50 | JN098762 | JN098882 | JN099120 | JN099163 | JN099001 | Theobroma cacoa | Indonesia | K.B. Boedijn \& J. Reitsma |
|  | CBS 114465=CPC 1619 | JN098763 | JN098887 | JN099125 | JN099170 | JN099008 |  | Ecuador | M.J. Wingfield |
| C. kurandica | CBS 129576=CPC 17547 | JN098764 | JN098941 | JN100634 | JN099233 | JN099071 | soil | Australia | P.W. Crous |
|  | CBS 129577= CPC 17551 ${ }^{\text {T }}$ | JN098765 | JN098953 | JN100646 | JN099245 | JN099083 | soil | Australia | P.W. Crous |
|  | CPC 17548 | JN098766 | JN098872 | JN099109 | JN099149 | JN098983 | soil | Australia | P.W. Crous |
|  | CPC 17549 | JN098767 | JN098871 | JN099108 | JN099148 | JN098982 | soil | Australia | P.W. Crous |
|  | CPC 17550 | JN098768 | JN098870 | JN099107 | JN099147 | JN098981 | soil | Australia | P.W. Crous |
|  | CPC 17553 | JN098769 | JN098869 | JN099106 | JN099146 | JN098980 | soil | Australia | P.W. Crous |
| C. lageniformis | CBS 340.92=PPRI 4449=UFV $115{ }^{\text {T }}$ | AY793481 | AY793520 | AF220959 | JN099165 | JN099003 | Eucalyptus sp. | Brazil | A.C. Alfenas |
|  | CBS 111060=CPC 1240 | JN098770 | JN098918 | JN100611 | JN099208 | JN099046 | Eucalyptus sp. | South Africa | P.W. Crous |
|  | CBS 111061=CPC 1241 | JN098771 | JN098913 | JN100606 | JN099202 | JN099040 | Eucalyptus sp. | South Africa | P.W. Crous |
|  | CBS 112898=CPC 5607 | AY725652 | AY725699 | AY793445 | JN099151 | JN098990 | Vitis vinifera | South Africa | L. Mostert |
| C. lageniformis | CBS 112899=CPC 5608 | AY793476 | AY793515 | AY793446 | JN099183 | JN099021 | Vitis vinifera | South Africa | L. Mostert |
|  | CBS 113011 $=$ CPC 4283 | JN098772 | JN098903 | JN100596 | JN099188 | JN099026 | Eucalyptus sp. | South Africa | P.W. Crous |
|  | CBS 113017=CPC 4287 | JN098773 | JN098884 | JN099122 | JN099167 | JN099005 | Eucalyptus sp. | South Africa | P.W. Crous |

Table 1 (continued)

| Species | Isolates | GenBank Accessions |  |  |  |  | Substrate | Country | Collector |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | BTUB | HIS3 | ITS | LSU | TEF-1 1 |  |  |  |
|  | CBS 113018=CPC 4286 | JN098774 | JN098904 | JN100597 | JN099189 | JN099027 | Eucalyptus sp. | South Africa | P.W. Crous |
|  | CBS 113019=CPC 4285 | JN098775 | JN098905 | JN100598 | JN099190 | JN099028 | Eucalyptus sp. | South Africa | P.W. Crous |
|  | CPC 17508 | JN098776 | JN098837 | JN100570 | JN099247 | JN099085 | soil | Australia | P.W. Crous |
|  | CPC 17509 | JN098777 | JN098935 | JN100628 | JN099227 | JN099065 | soil | Australia | P.W. Crous |
|  | CPC 17522 | JN098778 | JN098863 | JN099100 | JN099140 | JN098985 | soil | Australia | P.W. Crous |
|  | CPC 17523 | JN098779 | JN098948 | JN100641 | JN099240 | JN099078 | soil | Australia | P.W. Crous |
|  | CPC 17526 | JN098780 | JN098856 | JN099092 | JN099132 | JN098971 | soil | Australia | P.W. Crous |
|  | CPC 17527 | JN098781 | JN098947 | JN100640 | JN099239 | JN099077 | soil | Australia | P.W. Crous |
|  | CPC 17537 | JN098782 | JN098946 | JN100639 | JN099238 | JN099076 | soil | Australia | P.W. Crous |
|  | CPC 17540 | JN098783 | JN098926 | JN100619 | JN099217 | JN099055 | soil | Australia | P.W. Crous |
|  | CPC 17599 | JN098784 | JN098937 | JN100630 | JN099229 | JN099067 | soil | Australia | P.W. Crous |
|  | CPC 17600 | JN098785 | JN098938 | JN100631 | JN099230 | JN099068 | soil | Australia | P.W. Crous |
|  | CPC 18712 | JN098786 | JN098842 | JN100576 | JN099255 | JN098957 | Rosa sp. | USA | M. Munster |
| C. lanceolata | CBS 114950=CPC 396 | JN098787 | JN098898 | JN100591 | JN099181 | JN099019 | Eucalyptus sp. | South Africa | P.W. Crous |
|  | CBS 129565=CPC 17566 | JN098788 | JN098939 | JN100632 | JN099231 | JN099069 | soil | Australia | P.W. Crous |
|  | CBS 129566 $=$ CPC 17567 ${ }^{\text {T }}$ | JN098789 | JN098862 | JN099099 | JN099139 | JN098978 | soil | Australia | P.W. Crous |
| C. longiphialidica | CBS 129557=CPC 18839 ${ }^{\text {T }}$ | JN098790 | JN098851 | JN100585 | JN099264 | JN098966 | soil | Thailand | P.W. Crous |
|  | CBS 129558=CPC 18841 | JN098791 | JN098852 | JN100586 | JN099265 | JN098967 | soil | Thailand | P.W. Crous |
| C. longistipitata | CBS 112953=CPC 4720 | JN098792 | JN098902 | JN100595 | JN099187 | JN099025 | Opisthiolepsis heterophylla | Australia | C. Pearce \& B. Paulus |
|  | CBS 116075=CPC 708 ${ }^{\text {T }}$ | AY793506 | AY793546 | AF220958 | JN099155 | JN098993 | soil | China | M.J. Wingfield |
| C. microcylindrica | CBS 111794=ATCC 38571 = CPC 2375 ${ }^{\text {T }}$ | AY793483 | AY793523 | AY793452 | JN099203 | JN099041 | Echeveria elegans | Indonesia | C.F. Hill |
| C. natalensis | CBS 110800=CPC 529 | JN098793 | JN098915 | JN100608 | JN099205 | JN099043 | soil | South Africa | P.W. Crous |
|  | CBS 114943 $=$ CPC $456^{\text {T }}$ | JN098794 | JN098895 | JN100588 | JN099178 | JN099016 | Arachis hypogaea | South Africa | M.J. Wingfield |
|  | CBS 114944=CPC 457 | JN098795 | JN098896 | JN100589 | JN099179 | JN099017 | Arachis hypogaea | South Africa | M.J. Wingfield |
|  | CBS 114945=CPC 459 | JN098796 | JN098897 | JN100590 | JN099180 | JN099018 | Arachis hypogaea | South Africa | M.J. Wingfield |
|  | CPC 17395 | JN098797 | JN098936 | JN100629 | JN099228 | JN099066 | soil | Australia | P.W. Crous |
| C. nederlandica | CBS 143.95=PD94/1353 | JN098798 | JN098891 | JN099129 | JN099175 | JN099013 | Kalanchoë sp. | Netherlands | J.W. Veenbaas-Rijks |
|  | CBS 146.94=PD39/1776 | JN098799 | JN098889 | JN099127 | JN099173 | JN099011 | Rhododendron sp. | Netherlands |  |
|  | CBS 152.91=PD90/2015 ${ }^{\text {T }}$ | JN098800 | JN098910 | JN100603 | JN099195 | JN099033 | Pelargonium sp. | Netherlands | J.W. Veenbaas-Rijks |
| C. novaezelandica | CBS 486.77 $=$ ATCC $44815=$ CPC $2397{ }^{\text {T }}$ | AY793485 | AY793525 | AF220963 | JN099212 | JN099050 | Rhododendron indicum | New Zealand | H.J. Boesewinkel |
| C. parva | CBS 114524 $=$ ATCC 28272 $=$ CPC 2370 ${ }^{\text {T }}$ | AY793486 | AY793526 | AF220964 | JN099171 | JN099009 | Telopea speciosissima | New Zealand | H.J. Boesewinkel |
| C. peruviana | CBS 113022=CPC 4291 | JN098801 | JN098906 | JN100599 | JN099191 | JN099029 | Eucalyptus sp. | South Africa | P.W. Crous |
|  | CBS 114697=CPC 2573 | JN098802 | JN098886 | JN099124 | JN099169 | JN099007 | Vitis vinifera | South Africa | S. Lambrecht |
|  | CBS 114942=CPC 267 | JN098803 | JN098893 | JN100587 | JN099177 | JN099015 | Acacia mearnsii | South Africa | P.W. Crous |
|  | CBS 114952=CPC 398 | JN098804 | JN098854 | JN100572 | JN099251 | JN099089 | Eucalyptus sp. | South Africa | P.W. Crous |

Table 1 (continued)

| Species | Isolates | GenBank Accessions |  |  |  |  | Substrate | Country | Collector |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | BTUB | HIS3 | ITS | LSU | TEF-1 $\alpha$ |  |  |  |
| C. pseudocamelliae | CBS 114953 = CPC 399 | JN098805 | JN098885 | JN099123 | JN099168 | JN099006 | Eucalyptus sp. | South Africa | P.W. Crous |
|  | CBS 116089 $=$ CPC $640=$ UFO 200 | JN098806 | JN098875 | JN099112 | JN099154 | JN098969 | Piptadenia sp. | Brazil | A.O. Carvalho |
|  | CBS 116103=CPC 637=UFO 197 | JN098807 | JN098908 | JN100601 | JN099193 | JN099031 | Psidium guajava | Brazil | A.O. Carvalho |
|  | CPC 17517 | JN098808 | JN098944 | JN100637 | JN099236 | JN099074 | soil | Australia | P.W. Crous |
|  | CPC 17532 | JN098809 | JN098855 | JN099091 | JN099131 | JN098970 | soil | Australia | P.W. Crous |
|  | CPC 17533 | JN098810 | JN098940 | JN100633 | JN099232 | JN099070 | soil | Australia | P.W. Crous |
|  | CPC 17534 | JN098811 | JN098873 | JN099110 | JN099150 | JN098984 | soil | Australia | P.W. Crous |
|  | CPC 17535 | JN098812 | JN098945 | JN100638 | JN099237 | JN099075 | soil | Australia | P.W. Crous |
|  | CPC 17556 | JN098813 | JN098954 | JN100569 | JN099246 | JN099084 | soil | Australia | P.W. Crous |
|  | IMUR 1843 $=$ CPC $2404{ }^{\text {T }}$ | AY793500 | AY793540 | AF220966 | JN099266 | JN098968 | ants | Peru | M.P. Herrera |
|  | CBS 129555= CPC 18825 ${ }^{\text {T }}$ | JN098814 | JN098843 | JN100577 | JN099256 | JN098958 | soil | Thailand | P.W. Crous |
|  | CBS 129556=CPC 18832 | JN098815 | JN098846 | JN100580 | JN099259 | JN098961 | soil | Thailand | P.W. Crous |
|  | CPC 18826 | JN098816 | JN098844 | JN100578 | JN099257 | JN098959 | soil | Thailand | P.W. Crous |
|  | CPC 18836 | JN098817 | JN098849 | JN100583 | JN099262 | JN098964 | soil | Thailand | P.W. Crous |
|  | CPC 18838 | JN098818 | JN098850 | JN100584 | JN099263 | JN098965 | soil | Thailand | P.W. Crous |
| C. pseudohawaiiensis | CBS 210.94=PPRI 4450=UFV $125^{\text {T }}$ | JN098819 | JN098890 | JN099128 | JN099174 | JN099012 | Eucalyptus sp. | Brazil | A.C. Alfenas |
|  | CBS 115610=CPC 909 | JN098820 | JN098901 | JN100594 | JN099186 | JN099024 |  | Madagascar | P.W. Crous |
| C. pseudoinfestans | CBS 114530=CPC 2320 | JN098821 | JN098888 | JN099126 | JN099172 | JN099010 | soil | Madagascar | J.E. Taylor |
|  | CBS 114531 $=$ CPC $2319{ }^{\text {T }}$ | AY793508 | AY793548 | AF220957 | JN099166 | JN099004 | soil | Madagascar | J.E. Taylor |
| C. pseudoparva | CBS 113624=CPC 752 | JN098822 | JN098883 | JN099121 | JN099164 | JN099002 | Quercus sp | Switzerland | L. Petrini |
|  | CBS 122594 | JN098823 | JN098907 | JN100600 | JN099192 | JN099030 | Vitis riparia | New Zealand | K. Paice |
|  | CBS 129560=CPC 18149 ${ }^{\text {T }}$ | JN098824 | JN098927 | JN100620 | JN099218 | JN099056 | soil | Netherlands | P.W. Crous |
|  | CPC 18150 | JN098825 | JN098864 | JN099101 | JN099141 | JN098986 | soil | Netherlands | P.W. Crous |
| C. queenslandica | CBS 129574=CPC 17568 ${ }^{\text {T }}$ | JN098826 | JN098861 | JN099098 | JN099138 | JN098977 | soil | Australia | P.W. Crous |
|  | CBS 129575=CPC 17569 | JN098827 | JN098860 | JN099097 | JN099137 | JN098976 | soil | Australia | P.W. Crous |
| C. stellenboschensis | CBS 386.67 | JN098828 | JN098920 | JN100613 | JN099210 | JN099048 | Fragaria sp. | Netherlands | G.H. Boerema |
|  | CBS 110668= ${ }^{\text {CPC } 517}{ }^{\text {T }}$ | JN098829 | JN098922 | JN100615 | JN099213 | JN099051 | soil | South Africa | P.W. Crous |
|  | CBS 115611 $=$ CPC 4074 | JN098830 | JN098900 | JN100593 | JN099185 | JN099023 | Geum sp . | New Zealand | P.W. Crous |
|  | CBS 116170=CPC 753 | JN098831 | JN098894 | JN099117 | JN099160 | JN098998 | Quercus sp | Switzerland | L. Petrini |
|  | CBS 129559=CPC 15200 | JN098832 | JN098868 | JN099105 | JN099145 | JN098979 | leaf litter | Canada | P.W. Crous |
| C. thailandica | CBS 129570=CPC 18834 | JN098833 | JN098847 | JN100581 | JN099260 | JN098962 | soil | Thailand | P.W. Crous |
|  | CBS 129571 $=$ CPC 18835 ${ }^{\text {T }}$ | JN098834 | JN098848 | JN100582 | JN099261 | JN098963 | soil | Thailand | P.W. Crous |
|  | CPC 18831 | JN098835 | JN098845 | JN100579 | JN099258 | JN098960 | soil | Thailand | P.W. Crous |
| C. variabilis | CBS 375.93=IMI 317057 | JN098836 | JN098881 | JN099119 | JN099162 | JN099000 | Mangifera indica | India | P.N. Chowdhry |
|  | CBS 129561 $=$ CPC 17505 ${ }^{\text {T }}$ | JN098719 | JN098950 | JN100643 | JN099242 | JN099080 | soil | Australia | P.W. Crous |

Table 1 (continued)

| Species | Isolates | GenBank Accessions |  |  |  |  | Substrate | Country | Collector |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | BTUB | HIS3 | ITS | LSU | TEF-1 $\alpha$ |  |  |  |
| C. viticola | CBS 129562 $=$ CPC17506 | JN098720 | JN098951 | JN100644 | JN099243 | JN099081 | soil | Australia | P.W. Crous |
|  | CPC 17504 | JN098721 | JN098949 | JN100642 | JN099241 | JN099079 | soil | Australia | P.W. Crous |
|  | CPC 17563 | JN098722 | JN098933 | JN100625 | JN099223 | JN099061 | soil | Australia | P.W. Crous |
|  | CBS 112897 $=$ CPC 5606 ${ }^{\text {T }}$ | AY793504 | AY793544 | AY793468 | JN099226 | JN099064 | Vitis vinifera | South Africa | G.J. van Coller |
|  | CBS 114682=IMI 297470 $=$ CPC 2509 | JN098723 | JN098919 | JN100612 | JN099209 | JN099047 | Amorphophallus sp. | Thailand | R. Stevenson |
| Cylindrocladiella sp. | CBS 139.26 | JN098724 | JN098912 | JN100605 | JN099197 | JN099035 |  | Netherlands | C.J. Buisman |
|  | CBS 114960=CPC 375 | JN098725 | JN098874 | JN099111 | JN099152 | JN098991 | Pinus radiata | South Africa | P.W. Crous |
|  | CBS 114961= CPC 377 | JN098726 | JN098934 | JN100626 | JN099224 | JN099062 | Pinus radiata | South Africa | P.W. Crous |
|  | CBS 115687 $=$ CPC 530 | JN098727 | JN098909 | JN100602 | JN099194 | JN099032 | leaf litter | South Africa | P.W. Crous |
|  | CBS 115895=CPC 502 | JN098728 | JN098876 | JN099113 | JN099156 | JN098994 |  | South Africa | S. Lambrecht |
|  | CPC 374 | JN098729 | JN098841 | JN100575 | JN099254 | JN098956 | Pinus radiata | South Africa | P.W. Crous |
| Cylindrocladiella sp. | CBS 199.62 | JN098730 | JN098911 | JN100604 | JN099196 | JN099034 | Viburnum sp. | Netherlands | G.H. Boerema |
|  | CBS 110669 = CPC 509 | JN098731 | JN098914 | JN100607 | JN099204 | JN099042 | soil | South Africa | P.W. Crous |
| Cylindrocladiella sp. | CBS 874.68=ATCC 16315=IMI 299377 | JN098732 | JN098921 | JN100614 | JN099211 | JN099049 | soil | Germany | W. Gams |
| Cylindrocladiella sp. | CBS 100283 | JN098733 | JN098892 | JN099130 | JN099176 | JN099014 | twig on ground | Japan | H.-J. Schroers |
| Cylindrocladiella sp . | CBS 110946=CPC 970 | JN098734 | JN098917 | JN100610 | JN099207 | JN099045 |  |  | P.W. Crous |
|  | $\text { CBS } 115673=\text { CPC } 917$ | AY793502 | AY793542 | AY793466 | JN099153 | JN098992 | soil | South America | P.W. Crous |
|  | CBS 115675=CPC 968 | AY793503 | AY793543 | AY793467 | JN099184 | JN099022 | soil | South America | P.W. Crous |
| Cylindrocladiella sp. | CBS 112364 | AY793507 | AY793547 | AY793470 | JN099200 | JN099038 | Archontophoenix purpurea | Australia | F. Hill |
| Cylindrocladiella sp . | CBS 114780=CPC 278 | JN098735 | JN098925 | JN100618 | JN099216 | JN099054 | Eucalyptus grandis | South Africa | P.W. Crous |
|  | CBS 114884 $=$ CPC 279 | JN098736 | JN098924 | JN100617 | JN099215 | JN099053 | Eucalyptus grandis | South Africa | P.W. Crous |
| Cylindrocladiella sp. | CBS 114881 $=$ CPC 238 | JN098737 | JN098880 | JN099118 | JN099161 | JN098999 | Eucalyptus grandis | South Africa | P.W. Crous |
| Cylindrocladiella sp . | CBS 114885=CPC 262 | JN098738 | JN098923 | JN100616 | JN099214 | JN099052 | Eucalyptus sp. | South Africa | P.W. Crous |
| Cylindrocladiella sp. | CBS 114890= CPC 259 | JN098739 | JN098928 | JN100627 | JN099225 | JN099063 | Eucalyptus sp. | South Africa | P.W. Crous |
| Cylindrocladiella sp . | CBS 114957=CPC 426 | JN098740 | JN098899 | JN100592 | JN099182 | JN099020 | Eucalyptus sp. | South Africa | P.W. Crous |
|  | CPC 260 | JN098741 | JN098840 | JN100574 | JN099253 | JN098955 | Eucalyptus sp. | South Africa | P.W. Crous |
| Cylindrocladiella sp. | CBS 116095=CPC 678 | JN098742 | JN098879 | JN099116 | JN099159 | JN098997 | soil | South Africa | M.J. Wingfield |
| Cylindrocladiella sp . | CBS 122595 | JN098743 | JN098877 | JN099114 | JN099157 | JN098995 | Vitis riparia | New Zealand | K. Paice |
| Cylindrocladiella sp. | CPC 15198 | JN098744 | JN098930 | JN100622 | JN099220 | JN099058 | soil | Canada | P.W. Crous |
| Cylindrocladiella sp. | CPC 15199 | JN098745 | JN098952 | JN100645 | JN099244 | JN099082 | soil | Canada | P.W. Crous |
| Cylindrocladiella sp. | CPC 17603 | JN098746 | JN098865 | JN0990102 | JN099142 | JN098987 | soil | Australia | P.W. Crous |



 RNA, TEF-1 $\alpha=$ Translation elongation factor 1-alpha. ${ }^{\mathrm{T}}$ Ex-type cultures.
of the fungal structures mounted in $85 \%$ lactic acid. The conidial measurements are presented as the $95 \%$ confidence level with extremes in parentheses. Only the extremes are presented for other structures. The colony colours were determined on $2 \% \mathrm{w} / \mathrm{v}$ malt extract agar (MEA) after 7 day incubation at $24^{\circ} \mathrm{C}$ in the dark using the colour charts of Rayner (1970). Descriptions, nomenclature and illustrations were deposited in MycoBank (Crous et al. 2004).

## Results

## Phylogeny

Amplicons of approximately 530 bases were determined for BTUB, HIS3 and TEF- $1 \alpha$, 500 for ITS, and 850 for LSU. The phylogenetic analysis included 136 ingroup taxa, with Ca. pauciramosa (CBS 114861) and Ca. brachiatica (CBS 123700) as outgroup taxa. The initial NJ analysis of the LSU sequence data revealed that all the isolates included in the study formed a monophyletic clade (results not shown). Comparisons of the $70 \%$ reciprocal bootstrap NJ tree topologies of the individual gene regions showed no conflict and therefore the sequence datasets were combined. The resulting dataset of 2,956 characters, including alignment gaps, consisted of 2,070 constant and 131 parsimonyuninformative characters. Analysis of the 755 parsimonyinformative characters yielded 1,224 trees ( $\mathrm{TL}=, 4486$; $\mathrm{CI}=$ $0.308 ; \mathrm{RI}=0.843$; $\mathrm{RC}=0.260$ ), of which the first tree is presented (Fig. 1). For the Bayesian analysis, a HKY + I + G model was selected for BTUB and TEF-1 $\alpha$, GTR + I + G for HIS3 and LSU, and SYM + I + G for ITS which was incorporated into the analysis. The Bayesian consensus tree confirmed both the tree topology and bootstrap support of the strict consensus tree obtained with maximum-parsimony.

In the phylogenetic tree (Fig. 1) the Cylindrocladiella isolates are divided into two main clades. The first main clade [bootstrap support $(\mathrm{BS})=98$; posterior probability $(\mathrm{PP})=0.70]$ is further divided into two subclades. The first subclade $(\mathrm{BS}=98 ; \mathrm{PP}=0.70)$ represents C. novaezelandiae (CBS 486.77), C. elegans (CBS 338.92) and other closely related isolates that could represent novel phylogenetic species. The second subclade $(\mathrm{BS}=100 ; \mathrm{PP}=0.75)$ representing C. camelliae (CPC 234; Crous 2002) and C. peruviana (IMUR 1843) also consists of closely related isolates clustering together in smaller well-supported terminal clades, each representing possible novel species.

The second main clade ( $\mathrm{BS}=59 ; \mathrm{PP}=0.53$ ) is also divided into two subclades. In the first subclade $(\mathrm{BS}=82 ; \mathrm{PP}=$ 0.90 ) representing $C$. parva (CBS 114524) several isolates form well-supported terminal clades, also indicating potentially new species. The second subclade $(\mathrm{BS}=91 ; \mathrm{PP}=$ 0.93 ) further divides into a clade $(B S=89 ; \mathrm{PP}=0.65)$
representing $C$. viticola (CBS 112897), and a clade ( $\mathrm{BS}=100 ; \mathrm{PP}=1.00$ ) containing C. lageniformis (CBS 340.92 ) and $C$. infestans. This clade also consists of several well-supported terminal clades that could represent novel species. The ex-type strain of the anamorph state of C. infestans (CBS 111795) clustered ( $\mathrm{BS}=83$; $\mathrm{PP}=0.97$ ) separately from the ex-type strain of the purported teleomorph state of this species (CBS 114531, $\mathrm{BS}=100 ; \mathrm{PP}=0.98$ ), indicating that each ex-type strain represents a distinct species.

## Taxonomy

Based on the phylogenetic inference and morphological observations, numerous Cylindrocladiella isolates included in this study represent novel species. Following the approach of Lombard et al. (2009, 2010a-c) and Crous et al. (2006, 2008 , 2009) for other fungal groups, all new species are described in Cylindrocladiella, as this represents the older generic (Boesewinkel 1982), and best established name for this group of fungi.

Cylindrocladiella australiensis L. Lombard \& Crous, sp. nov. - MycoBank MB561676, Fig. 2.

Etymology - Named after the country from where it was collected, Australia.

Cylindrocladiellae infestantis morphologice valde similis, sed conidiis minoribus, $(9-) 11-13(-15) \times 2-4 \mu \mathrm{~m}$, distinguitur.

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 2a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, $41-96 \times 6-9 \mu \mathrm{~m}$; stipe extension aseptate, straight, $101-152 \mu \mathrm{~m}$ long, thick-walled with one basal septum, terminating in thin-walled, ellipsoidal to fusoid vesicles (Fig. 2j-1), 6-8 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig. 2f-i) with primary branches aseptate, 13-21× $3-5 \mu \mathrm{~m}$, secondary branches aseptate, $11-15 \times 3-6 \mu \mathrm{~m}$, each terminal branch producing $2-4$ phialides; phialides cymbiform to cylindrical, hyaline, aseptate, $8-17 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores (Fig. 2m-n) abundant, comprising of a septate stipe, primary and secondary branches terminating in 2-3 phialides; primary branches straight, hyaline, $0-1$-septate, $22-54 \times 2-5 \mu \mathrm{~m}$, secondary branches straight, hyaline, aseptate, $21-36 \times 4-5 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $19-40 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 2o) cylindrical, rounded at both ends, straight, 1 -septate, (9-)11-13(-15) $\times 2-4 \mu \mathrm{~m}$ (av. $=12 \times 3 \mu \mathrm{~m}$ ), frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.


Fig. 1 One of 1,224 most parsimonious trees obtained from a heuristic search with 1,000 random additions sequences of the combined $\beta$-tubulin, histone H3, internal transcribed spacer regions 1 and 2 and the 5.8 S gene of the ribosomal RNA, 28S large subunit and translation elongation factor-1alpha sequence alignments of the Cylindrocladiella isolates. Scale bar shows 10 changes and bootstrap support values (bold) from 1,000 replicates
and Bayesian posterior probability values are indicated at the nodes. Thickened lines indicate branches in the strict consensus tree and the consensus tree of the Bayesian analyses. Ex-type strains are indicated in bold and coloured block indicate the novel species described. The tree was rooted to Calonectria brachiatica (CBS123700) and Ca. pauciramosa (CBS 114861). Species complexes are indicated on the right

$-{ }^{10}$
Fig. 1 (continued)


Fig. 2 Cylindrocladiella australiensis. a-e. Penicillate conidiophores. f-i. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{l}$. Terminal vesicles. $\mathbf{m}-\mathbf{n}$. Subverticillate conidiophores. o. Conidia. $A=50 \mu \mathrm{~m}($ apply to $\mathbf{b}-\mathbf{e}, \mathbf{m}), \mathrm{F}=10 \mu \mathrm{~m}($ apply to $\mathbf{g}-\mathbf{l}, \mathbf{n}-\mathbf{o})$

Culture characteristics - Colonies raised (convex), cottony, with smooth margins, white, buff yellow (19 d) to umber (13i) (reverse); chlamydospores moderate throughout medium, arranged in chains; reaching 90 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - Australia, Queensland, Daydream Island, Whitsundays Island Resort, from soil, 2 Aug. 2009, P.W. Crous, holotype CBS-H20596, culture ex-type CBS $129567=$ CPC 17507; Australia, Queensland, Lake Barrine,
from soil, 18 June 2009, P.W. Crous, culture CBS $129568=$ CPC 17562.

Notes - Cylindrocladiella australiensis can be distinguished from C. infestans (av. $15 \times 3 \mu \mathrm{~m}$ ) by its smaller conidia and its terminal vesicle shape. The subverticillate conidiophores of C. australiensis also form secondary branches not reported for C. infestans. Unique fixed nucleotides were also identified for C. australiensis for three loci: BTUB positions 186 (T), 296 (C), 350 (T),

381 (C) and $387(\mathrm{~T})$; HIS3 positions $90(\mathrm{~T})$ and $387(\mathrm{~T})$; TEF$1 \alpha$ positions $113(\mathrm{C}), 153(\mathrm{G}), 155(\mathrm{C}), 168(\mathrm{~T}), 229(\mathrm{~T}), 232$ (C), 254 (G), 266 (C), 282 (T), 462 (A) and 468 (C).

Cylindrocladiella clavata L. Lombard \& Crous, sp. nov. MycoBank MB561674, Fig. 3.

Etymology - Named after the clavate shape of its vesicles.

Cylindrocladiellae variabilis morphologice similis, sed vesiculis clavatis distinguitur.

Teleomorph unknown. Conidiophores monomorphic, penicillate, mononematous and hyaline. Penicillate conidiophores (Fig. 3a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, $40-86 \times 6-$ $10 \mu \mathrm{~m}$; stipe extension aseptate, straight, 116-170 $\mu \mathrm{m}$ long, thick-walled with one basal septum, terminating in thin-walled, elongated, clavate vesicles (Fig. 3f-h), 4-7 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig. 3i-k) with primary branches aseptate, $10-23 \times 3-8 \mu \mathrm{~m}$, secondary branches
aseptate, $6-11 \times 2-4 \mu \mathrm{~m}$, each terminal branch producing 2-4 phialides; phialides doliiform to cymbiform, hyaline, aseptate, $7-12 \times 2-3 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette absent. Subverticillate conidiophores not observed. Conidia (Fig. 31) cylindrical, rounded at both ends, straight, 1 -septate, (10-)13-15($16) \times 2-3 \mu \mathrm{~m}(\mathrm{av} .=14 \times 2 \mu \mathrm{~m})$, frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with undulate margins, white with buff yellow (19 d) centre, umber (13i) (reverse); chlamydospores extensive throughout medium arranged in chains; reaching 70 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimen examined - Australia, Queensland, Byron Bay, from soil, 17 July 2009, P.W. Crous, holotype CBS-H20597, culture ex-type CBS 129564=CPC 17592, Australia, Queensland, Byron Bay, from soil, 17 July 2009, P.W. Crous, culture CBS $129563=$ CPC 17591.


Fig. 3 Cylindrocladiella clavata. a-e. Penicillate conidiophores. $\mathbf{f}-\mathbf{h}$. Terminal vesicles. $\mathbf{i}-\mathbf{k}$. Conidiogenous apparatus with conidiophore branches and phialides. L. Conidia. Scale bars: $\mathrm{A}=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{e}), \mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{g - l}$ )

Notes - This species can be distinguished from other species in the genus by its elongated clavate terminal vesicles. The conidia are also slightly larger than those of C. lageniformis (av. $12 \times 2 \mu \mathrm{~m}$ ).

Cylindrocladiella cymbiformis L. Lombard \& Crous, sp. nov. - MycoBank MB561666, Fig. 4.

Etymology - Named after its phialides, which are cymbiform in shape.

Cylindrocladiellae elegantis morphologice valde similis, sed conidis majoribus, $(15-) 16-20(-22) \times(2-) 3-5(-6) \mu \mathrm{m}$, distinguitur.

Teleomorph unknown. Conidiophores monomorphic, penicillate, mononematous, hyaline; comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle (Fig. 4a-e); stipe septate, hyaline, smooth, $44-84 \times 2-4 \mu \mathrm{~m}$; stipe extension aseptate, straight, $107-175 \mu \mathrm{~m}$ long, thick-walled with one basal septum, terminating in thin-walled, lageniform to broadly clavate
vesicles (Fig. 4j-1), 6-8 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig. 4f-i) with primary branches aseptate, $12-$ $28 \times 4-6 \mu \mathrm{~m}$, secondary branches aseptate, $11-19 \times 2-5 \mu \mathrm{~m}$, each terminal branch producing $2-4$ phialides; phialides cymbiform, hyaline, aseptate, $11-19 \times 2-5 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores not observed. Conidia (Fig. 4m) cylindrical, rounded at both ends, straight, 1 -septate, (15-)16-20($22) \times(2-) 3-5(-6) \mu \mathrm{m}(\mathrm{av} .=18 \times 3 \mu \mathrm{~m})$, frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with undulate margins, white, buff yellow (19 d) (reverse); chlamydospores sparse throughout medium, arranged in chains; reaching 45 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimen examined - Australia, Queensland, Brisbane, from soil, 11 July 2009, P.W. Crous, holotype CBS-H20598,


Fig. 4 Cylindrocladiella cymbiformis. a-e. Penicillate conidiophores. f-i. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{l}$. Terminal vesicles. $\mathbf{m}$. Conidia. Scale bars: $A=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{e}), \mathrm{F}=10 \mu \mathrm{~m}($ apply to $\mathbf{g}-\mathbf{m})$
culture ex-type CBS 129553=CPC 17393; Australia, Queensland, Brisbane, from soil, 11 July 2009, P.W. Crous, culture CBS 129554=CPC 17392.

Notes - Based on phylogenetic inference, C. cymbiformis is placed in the C. elegans species complex, and closely related to C. novaezelandiae. Morphologically, this species has larger conidia (av. $18 \times 3 \mu \mathrm{~m}$ ) than C. elegans (av. $14.5 \times$ $2 \mu \mathrm{~m}$ ) and C. novaezelandiae (av. $14.5 \times 2 \mu \mathrm{~m}$ ) (Crous 2002), and stipe extensions are also much longer. Only cymbiform phialides were observed for C. cymbiformis, whereas both C. elegans and C. novaezelandiae also produce doliiform to reniform phialides. Furthermore, no subverticillate conidiophores were observed for C. cymbiformis, but have been reported for C. elegans.

Cylindrocladiella ellipsoidea L. Lombard \& Crous, sp. nov. - MycoBank MB561681, Fig. 5.

Etymology - Named after the characteristic ellipsoid shape of its vesicles.

Cylindrocladiellae infestantis morphologice similis, sed vesiculis clavatis vel ellipsoideis distiunguitur.

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 5a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, 41$83 \times 6-8 \mu \mathrm{~m}$; stipe extension aseptate, straight, $77-155 \mu \mathrm{~m}$ long, thick-walled with one basal septum, terminating in thin-walled, clavate to ellipsoidal vesicles (Fig. 5j-n), 5$8 \mu \mathrm{~m}$ wide. Penicillate conidiogenous apparatus (Fig. 5f-i) with primary branches aseptate, $11-20 \times 3-6 \mu \mathrm{~m}$, secondary branches aseptate, $9-12 \times 3-5 \mu \mathrm{~m}$, each terminal branch producing $2-4$ phialides; phialides doliiform to cymbiform, hyaline, aseptate, $9-12 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores (Fig. $50-\mathrm{p}$ ) in moderate numbers, comprising of a septate stipe, primary and secondary branches terminating in $2-3$ phialides; primary branches straight, hyaline, $0-$ 1 -septate, $23-40 \times 3-6 \mu \mathrm{~m}$, secondary branches rare, straight, hyaline, aseptate, $16-31 \times 4 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $27-52 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 5q) cylindrical, rounded at both ends, straight, 1-septate, (14-)16-$18(-19) \times 3-4 \mu \mathrm{~m}(\mathrm{av} .=17 \times 4 \mu \mathrm{~m})$, frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth to undulate margins, white, buff yellow (19 d) to umber (13i) (reverse); chlamydospores moderate throughout media arranged in chains; reaching 60 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimen examined - Australia, Queensland, Lake Barrine, from soil, 18 June 2009, P.W. Crous, holotype CBS-H20599, culture ex-type CBS 129573=CPC 17560;

Australia, Queensland, Lake Barrine, from soil, 18 June 2009, P.W. Crous, culture CBS 129572=CPC 17558; Australia, Queensland, Lake Barrine, from soil, 18 June 2009, P.W. Crous, culture CPC 17559.

Notes - Cylindrocladiella ellipsoidea produces subverticillate conidiophores with secondary branches, which has not reported been observed for other species in the $C$. infestans complex. Furthermore, C. ellipsoidea can also be distinguished for others in the complex based on their terminal vesicle shape. Unique fixed nucleotides were also identified for $C$. ellipsoidea for two loci: HIS3 positions 124 (C), 130 (A), 134 (C), 314 (A) and 349 (A); TEF-1 $\alpha$ position 210 (indel).

Cylindrocladiella hawaiiensis L. Lombard \& Crous, sp. nov. - MycoBank MB561677, Fig. 6.

Etymology - Named after Hawaii, where this fungus was collected.

Cylindrocladiellae infestantis morphologice similis, sed vesiculis clavatis distinguitur.

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 6a-c) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, 47$80 \times 5-6 \mu \mathrm{~m}$; stipe extension aseptate, straight, $80-116 \mu \mathrm{~m}$ long, thick-walled with one basal septum, terminating in thin-walled, clavate vesicles (Fig. 6d-e), 5-7 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig. $6 \mathrm{f}-\mathrm{j}$ ) with primary branches aseptate, $11-19 \times 4-5 \mu \mathrm{~m}$, secondary branches aseptate, $8-19 \times 3-4 \mu \mathrm{~m}$, each terminal branch producing $2-$ 4 phialides; phialides cymbiform to cylindrical, hyaline, aseptate, $8-18 \times 2-3 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores (Fig. 6k-1) abundant, comprising of a septate stipe, and primary branches terminating in 2-3 phialides; primary branches straight, hyaline, $0-1$-septate, $23-38 \times 3-5 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $19-41 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 6m) cylindrical, rounded at both ends, straight, 1septate, $(10-) 12-14 \times 2-4 \mu \mathrm{~m}(\mathrm{av} .=13 \times 3 \mu \mathrm{~m})$, frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth margins, white, buff yellow (19 d) (reverse); chlamydospores sparse throughout medium, arranged in chains; reaching 65 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - Hawaii, from soil, 1 Aug. 2005, Y. Degawa, holotype CBS-H20600, culture ex-type CBS $129569=$ CPC 12272; Hawaii, Kaua'i Island, Secret waterfall, from soil, 8 Aug. 2005, Y. Degawa, culture CBS 118704.

Notes - Cylindrocladiella hawaiiensis produces clavate terminal vesicles, distinguishing it from C. infestans, which


Fig. 5 Cylindrocladiella ellipsoidea. a-e. Penicillate conidiophores. $\mathbf{f}-\mathbf{i}$. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{n}$. Terminal vesicles. o-p. Subverticillate conidiophores. $\mathbf{q}$.

Conidia. Scale bars: $\mathrm{A}=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{c}$ ), $\mathrm{D}=20 \mu \mathrm{~m}$ (apply to $\mathbf{e}$, o), $\mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{g - n}, \mathbf{p}-\mathbf{q}$ )
(A), 229 (A), 235 (G), 248 (A), 278 (A), 456 (indel), 462 (T) and 469 (A).

Cylindrocladiella kurandica L. Lombard \& Crous, sp. nov. - MycoBank MB561683, Fig. 7.

Etymology - Named after the Kuranda, the town where this fungus was collected.


Fig. 6 Cylindrocladiella hawaiiensis. a-c. Penicillate conidiophores. d-e. Terminal vesicles. $\mathbf{f}-\mathbf{j}$. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{k}-\mathbf{l}$. Subverticillate conidiophores. $\mathbf{m}$. Conidia. $\mathrm{A}=20 \mu \mathrm{~m}$ (apply to $\mathbf{b}, \mathbf{j}$ ), $\mathrm{D}=10 \mu \mathrm{~m}$ (apply to $\mathbf{e}-\mathbf{i}, \mathbf{k}-\mathbf{m}$ )

Cylindrocladiellae infestantis morphologice valde similis et vix distinguibilis, sed characteribus sequentibus nucleotiditis fixationibus in positionibus diversis [BTUB 97 (T), 395 (A) et 482 (T); HIS3 22 (T), 50 (A) et 315 (T); TEF$1 \propto 107(\mathrm{C})]$ genetice distinguitur.

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 7a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, $54-87 \times 5-9 \mu \mathrm{~m}$; stipe extension aseptate, straight, 153-219 $\mu \mathrm{m}$ long, thick-walled with one basal septum, terminating in thin-walled, ellipsoidal to lanceolate vesicles (Fig. 7j-1), 6-9 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig. $7 \mathrm{f}-\mathrm{i}$ ) with primary branches aseptate, $12-24 \times$ $3-7 \mu \mathrm{~m}$, secondary branches aseptate, $8-15 \times 2-4 \mu \mathrm{~m}$, each
terminal branch producing 2-4 phialides; phialides doliiform to cymbiform, hyaline, aseptate, $8-14 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores (Fig. 7m-o) in moderate numbers, comprising of a septate stipe, and primary branches terminating in 2-3 phialides; primary branches straight, hyaline, $0-1$-septate, $20-$ $48 \times 2-4 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $18-35 \times 2-5 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 7p) cylindrical, rounded at both ends, straight, 1 -septate, (10-)12-14($16) \times 2-4 \mu \mathrm{~m}(\mathrm{av} .=13 \times 3 \mu \mathrm{~m})$, frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised, cottony, with undulate margins, white with straw ( 21 d ) tint in patches, umber (13i) (reverse); chlamydospores extensive


Fig. 7 Cylindrocladiella kurandica. a-e. Penicillate conidiophores. fi. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{l}$. Terminal vesicles. $\mathbf{m}-\mathbf{0}$. Subverticillate conidiophores. p. Conidia.

Scale bars: $\mathrm{A}=50 \mu \mathrm{~m}, \mathrm{~B}=20 \mu \mathrm{~m}$ (apply to $\mathbf{c}-\mathbf{e}, \mathbf{m}-\mathbf{o}$ ), $\mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{g}-\mathrm{l}, \mathbf{p}$ )
throughout medium, arranged in chains; reaching 65 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimen examined - Australia, Queensland, Kuranda, from soil, 13 Aug 2009, P.W. Crous, holotype CBSH20601, culture ex-type CBS 129577=CPC 17551; Australia, Queensland, Kuranda, from soil, 13 Aug 2009, P.W. Crous, culture CBS 129576=CPC 17547; Australia,

Queensland, Kuranda, from soil, 13 Aug 2009, P.W. Crous, culture CPC 17549.

Notes - Cylindrocladiella kurandica is difficult to distinguish from C. longistipitata and other species in the C. infestans complex, and therefore phylogenetic inference is required for an accurate identification. Cylindrocladiella kurandica can be distinguished from other species in the C. infestans
complex by different unique fixed nucleotides for three loci: BTUB positions 97 (T), 395 (A) and 482 (T); HIS3 positions $22(\mathrm{~T}), 50(\mathrm{~A})$ and $315(\mathrm{~T})$; TEF- $1 \alpha$ position $107(\mathrm{C})$.

Cylindrocladiella lanceolata L. Lombard \& Crous, sp. nov. - MycoBank MB561675, Fig. 8.

Etymology - Named after the lanceolate shape of its vesicles.

Cylindrocladiellae lageniformis morphologice similis, sed vesiculis lanceolatis distinguitur.

Teleomorph unknown. Conidiophores monomorphic, penicillate, mononematous and hyaline. Penicillate conidiophores (Fig. 8a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, 31-77× $5-10 \mu \mathrm{~m}$; stipe extension aseptate, straight, 76-173 $\mu \mathrm{m}$ long, thick-walled with one basal septum, terminating in thin-walled, lanceolate vesicles (Fig. 8f-h), 5-7 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig. 8i-k) with primary branches aseptate, $12-30 \times 3-8 \mu \mathrm{~m}$,
secondary branches aseptate, $7-17 \times 3-6 \mu \mathrm{~m}$, each terminal branch producing 2-4 phialides; phialides reniform to doliiform to cymbiform, hyaline, aseptate, $7-13 \times 2-$ $3 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette absent. Subverticillate conidiophores not observed. Conidia (Fig. 81) cylindrical, rounded at both ends, straight, 1 -septate, $(13-) 15-17(-20) \times 2-3 \mu \mathrm{~m}$ (av. $=16 \times 3 \mu \mathrm{~m}$ ), frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth margins, white, umber (13i) (reverse); chlamydospores extensive throughout medium, arranged in chains; reaching 55 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - Australia, Queensland, Brisbane, from soil, 18 July 2009, P.W. Crous, holotype CBS-H20602, culture ex-type CBS 129566=CPC 17567, CBS 129565= CPC 17566; South Africa, KwaZulu-Natal, Kwambonambi, Mondi Sawmill, from Eucalyptus sp., 1 May 1990, P.W. Crous, culture CBS $114950=$ CPC 396.


Fig. 8 Cylindrocladiella lanceolata. a-e. Penicillate conidiophores. f-h. Terminal vesicles. i-k. Conidiogenous apparatus with conidiophore branches and phialides. L. Conidia. Scale bars: $\mathrm{A}=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{e}$ ), $\mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{g}-\mathbf{l}$ )

Note - Cylindrocladiella lanceolata can be distinguished from C. lageniformis by its lanceolate terminal vesicles and conidium dimensions.

Cylindrocladiella longiphialidica L. Lombard \& Crous, sp. nov. - MycoBank MB561669, Fig. 9.

Etymology - Named after its characteristically long phialides.

Cylindrocladiellae camelliae morphologice valde similis, sed phialidibus conidiophorum subverticillatorum longioribus distinguitur.


Fig. 9 Cylindrocladiella longiphialidica. a-e. Penicillate conidiophores. $\mathbf{f}-\mathbf{i}$. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{n}$. Terminal vesicles. o-q. Subverticillate conidiophores. R. Conidia. Scale bars: $A=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{e}$ ), $\mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{g}-\mathbf{r}$ )

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 9a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, 43$107 \times 6-9 \mu \mathrm{~m}$; stipe extension aseptate, straight, 114$189 \mu \mathrm{~m}$ long, thick-walled with one basal septum, terminating in thin-walled, ellipsoidal to lanceolate vesicles (Fig. 9j-n), 5$8 \mu \mathrm{~m}$ wide. Penicillate conidiogenous apparatus (Fig. 9f-i) with primary branches aseptate, $11-33 \times 3-7 \mu \mathrm{~m}$, secondary branches aseptate, $9-26 \times 3-5 \mu \mathrm{~m}$, with each terminal branch producing $2-4$ phialides; phialides doliiform to reniform to cymbiform, hyaline, aseptate, $8-13 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores (Fig. 90-q) abundant, comprising of a septate stipe, and primary branches terminating in 1-3 phialides; primary branches straight, hyaline, $0-1$-septate, $28-68 \times 4$ $6 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $20-79 \times 2-5 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 9r) cylindrical, rounded at both ends, straight, 1 -septate, $12-14 \times 2-3 \mu \mathrm{~m}$ (av. $=13 \times$ $3 \mu \mathrm{~m}$ ), frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with undulate margins, white centre becoming buff yellow (19 d) towards the margins, buff yellow (19 d) (reverse); chlamydospores extensive throughout medium, arranged in chains; reaching 55 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - Thailand, Chiang Mai, from soil, Oct. 2010, P.W. Crous, holotype CBS-H20603, culture ex-type CBS 129557=CPC 18839; Thailand, Chiang Mai, from soil, Oct. 2010, P.W. Crous culture, CBS 129558= CPC 18841.

Notes - Cylindrocladiella longiphialidica is morphologically similar to C. nederlandica, C. pseudocamelliae and C. camelliae, but can be distinguished from these species by its longer phialides on the subverticillate conidiophores.

Cylindrocladiella longistipitata L. Lombard \& Crous, sp. nov. - MycoBank MB561679, Fig. 10.

Etymology - Named after its characteristically long stipe extensions on its conidiophores.

Cylindrocladiellae infestantis morphologice similis, sed extensionibus stipitis longioribus.

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 10a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, $54-80 \times 5-9 \mu \mathrm{~m}$; stipe extension aseptate, straight, 130$216 \mu \mathrm{~m}$ long, thick-walled with one basal septum, terminating in thin-walled, cylindrical to lanceolate vesicles (Fig. 10j-n), $5-7 \mu \mathrm{~m}$ wide. Penicillate conidiogenous apparatus
(Fig. 10f-i) with primary branches aseptate, $13-20 \times 3-$ $5 \mu \mathrm{~m}$, secondary branches aseptate, $9-13 \times 3-5 \mu \mathrm{~m}$, each terminal branch producing 2-4 phialides; phialides cymbiform to cylindrical, hyaline, aseptate, $10-16 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores (Fig. 10o-p) in moderate numbers, comprising of a septate stipe, and primary branches terminating in 2-3 phialides; primary branches straight, hyaline, $0-1$-septate, $21-40 \times 4 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $18-31 \times 2-$ $4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 10q) cylindrical, rounded at both ends, straight, 1-septate, (12-)14-16(-17) $\times 2-4 \mu \mathrm{~m}$ (av. $=$ $15 \times 3 \mu \mathrm{~m}$ ), frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth to undulate margins, white, umber (13i) (reverse); chlamydospores extensive throughout medium, arranged in chains; reaching 45 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - China, Hong Kong, from soil, Nov. 1993, M.J. Wingfield, holotype CBS-H20604, culture ex-type CBS 116075=CPC 708; Australia, Queensland, Topaz, Atherton Tablelands, from Opisthiolepsis heterophylla, 2 Apr. 2001, C. Pearce \& B. Paulus, culture CBS $112953=$ CPC 4720.

Notes - Cylindrocladiella longistipitata can be distinguished from other species in the $C$. infestans complex by its longer stipe extension and terminal vesicle morphology. Furthermore, it has unique fixed nucleotides for three loci: BTUB position 363 (A); HIS3 positions 37 (C) and $400(\mathrm{~T})$; TEF-1 $\alpha$ positions 44 (A) and 45 (T).

Cylindrocladiella natalensis L. Lombard \& Crous, sp. nov. - MycoBank MB561670, Fig. 11.

Etymology - Named after the Province in South Africa where this fungus was first collected, KwaZulu-Natal.

Cylindrocladiellae elegantis morphologice valde similis, sed conidiis majoribus, $(12-) 14-16(-17) \times 2-3 \mu \mathrm{~m}$, distinguitur.

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 11a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, $88-135 \times 5-8 \mu \mathrm{~m}$; stipe extension aseptate, straight, $82-$ $127 \mu \mathrm{~m}$ long, thick-walled with one basal septum, terminating in thin-walled, ellipsoidal to fusoid vesicles (Fig. 11j-m), 6$8 \mu \mathrm{~m}$ wide. Penicillate conidiogenous apparatus (Fig. 11f-i) with primary branches aseptate, $13-29 \times 2-5 \mu \mathrm{~m}$, secondary branches aseptate, $8-17 \times 3-4 \mu \mathrm{~m}$, each terminal branch producing 2-4 phialides; phialides cymbiform to cylindrical, hyaline, aseptate, $9-14 \times 2-3 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores


Fig. 10 Cylindrocladiella longistipitata. a-e. Penicillate conidiophores. f-i. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{n}$. Terminal vesicles. $\mathbf{0}-\mathbf{p}$. Subverticillate conidiophores. Q. Conidia. $A=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{e}$ ), $\mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{g}-\mathbf{q}$ )
(Fig. 11n-o) in moderate numbers, comprising of a septate stipe, and primary branches terminating in 2-3 phialides; primary branches straight, hyaline, $0-1$-septate, $23-39 \times 2-$ $4 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $19-34 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 11p) cylindrical, rounded at both ends, straight, 1 -septate, (12-)14-16(-17) $\times 2-3 \mu \mathrm{~m}$ (av. $=$ $15 \times 3 \mu \mathrm{~m}$ ), frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth to undulate margins, white, buff yellow (19 d) (reverse); chlamydospores sparse throughout medium, arranged in chains; reaching 70 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - South Africa, KwaZulu-Natal, from Arachis hypogaea, 1 Feb. 1991, M.J. Wingfield, holotype CBS-H20605, culture ex-type CBS 114943= CPC 456, CBS 114945=CPC 459; Australia, Queensland,


Fig. 11 Cylindrocladiella natalensis. a-e. Penicillate conidiophores. f-i. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{m}$. Terminal vesicles. $\mathbf{n}-\mathbf{0}$. Subverticillate conidiophores. P. Conidia. Scale bars: $A=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{e}$ ), $\mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{g}-\mathbf{p}$ )

Byron Bay, from soil, 17 July 2009, P.W. Crous, culture CPC 17395.

Note - Cylindrocladiella natalensis can be distinguished from other species in this genus by its conidium dimensions and shape of the terminal vesicle.

Cylindrocladiella nederlandica L. Lombard \& Crous, sp. nov. - MycoBank MB561667, Fig. 12.

Etymology - Named after the Netherlands, the country where this fungus was collected.

Cylindrocladiellae camelliae morphologice valde similis, sed phialidibus majoribus, $14-30 \times 3-5 \mu \mathrm{~m}$, distinguitur.

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 12a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, $41-124 \times 4-10 \mu \mathrm{~m}$; stipe extension aseptate, straight, $102-$ $158 \mu \mathrm{~m}$ long, thick-walled with one basal septum,


Fig. 12 Cylindrocladiella nederlandica. a-e. Penicillate conidiophores. f-i. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{n}$. Terminal vesicles. $\mathbf{0}-\mathbf{p}$. Subverticillate conidiophores. Q. Conidia. Scale bars: $\mathrm{A}=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{e}$ ), $\mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{g}-\mathbf{q}$ )
terminating in thin-walled, lageniform to ellipsoidal vesicles (Fig. 12j-n), 4-9 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig. 12f-i) with primary branches aseptate, $12-31 \times 3-7 \mu \mathrm{~m}$, secondary branches aseptate, $8-18 \times 2-$ $5 \mu \mathrm{~m}$, each terminal branch producing $2-4$ phialides; phialides doliiform to reniform to cymbiform, hyaline, aseptate, $8-14 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal
thickening and collarette. Subverticillate conidiophores (Fig. 12o-p) abundant, comprising of a septate stipe, and primary branches terminating in 1-3 phialides; primary branches straight, hyaline, $0-1$-septate, $18-32 \times 3-5 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $14-30 \times 3-5 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 12q) cylindrical, rounded at
both ends, straight, 1 -septate, (10-)12-14(-15) $\times 2-4 \mu \mathrm{~m}$ $(\mathrm{av} .=13 \times 2 \mu \mathrm{~m})$, frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth to undulate margins, white, buff yellow (19 d) to umber (13i) (reverse); chlamydospores moderate throughout medium, arranged in chains; reaching 55 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - The Netherlands, from Pelargonium sp., Mar. 1991, J.W. Veenbaas-Rijks, holotype CBS-H5129, culture ex-type CBS $152.91=$ PD 90/2015; The Netherlands, Aalsmeer, from Kalanchoë sp., Feb. 1995, J.W. Veenbaas-Rijks, culture CBS $143.95=$ PD 94/1353; The Netherlands, stem of Rhododendron, Mar. 1994, culture CBS 146.94=PD 39/1776.

Notes - Morphologically, isolates of C. nederlandica are very similar to $C$. camelliae, with a slight difference in terminal vesicle shape. The phialides on the subverticillate conidiophores of $C$. nederlandica $(14-30 \times 3-5 \mu \mathrm{~m})$ are larger than those of $C$. camelliae $(15-26 \times 2-3.5 \mu \mathrm{~m}$; Crous 2002).

Cylindrocladiella pseudocamelliae L. Lombard \& Crous, sp. nov. - MycoBank MB561668, Fig. 13.

Etymology - Named after its morphological similarity to Cylindrocladiella camelliae.

Cylindrocladiellae camelliae morphologice similis, sed vesiculis divergentibus.

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 13a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, $65-137 \times 6-10 \mu \mathrm{~m}$; stipe extension aseptate, straight, $106-$ $188 \mu \mathrm{~m}$ long, thick-walled with one basal septum, terminating in thin-walled, ellipsoidal to lageniform to lanceolate vesicles (Fig. 13j-n), 6-10 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig. 13f-i) with primary branches aseptate, $12-27 \times 3-6 \mu \mathrm{~m}$, secondary branches aseptate, $8-18 \times 2-$ $5 \mu \mathrm{~m}$, each terminal branch producing $2-4$ phialides; phialides doliiform to reniform to cymbiform, hyaline, aseptate, $10-17 \times 2-3 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores (Fig. 13o-p) abundant, comprising of a septate stipe, and primary branches terminating in $1-3$ phialides; primary branches straight, hyaline, $0-1$-septate, $15-32 \times 3-6 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $19-31 \times 3-5 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 13q) cylindrical, rounded at both ends, straight, 1 -septate, (9-)11-15(-16) $\times 2-4 \mu \mathrm{~m}$ (av. $=13 \times 3 \mu \mathrm{~m}$ ), frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth margins, white, buff yellow (19 d) to umber (13i)
(reverse); chlamydospores moderate throughout medium, arranged in chains; reaching 90 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - Thailand, Chiang Mai, from soil, Oct. 2010, P.W. Crous, holotype CBS-H20606, culture ex-type CBS 129555=CPC 18825; Thailand, Chiang Mai, from soil, Oct. 2010, P.W. Crous, culture CBS 129556=CPC 18832; Thailand, Chiang Mai, from soil, Oct. 2010, P.W. Crous, culture CPC 18838.

Notes - As with C. nederlandica, C. pseudocamelliae is morphologically similar to C. camelliae. However, C. pseudocamelliae can be distinguished from both the other species by its longer stipe extension and the shape of its terminal vesicle.

Cylindrocladiella pseudohawaiiensis L. Lombard \& Crous, sp. nov. - MycoBank MB561678, Fig. 14.

Etymology - Named after its morphological similarity to Cylindrocladiella hawaiiensis.

Cylindrocladiellae infestantis morphologice valde similis, sed conidiis minoribus, (11-)12-14(-15) $\times 2-4 \mu \mathrm{~m}$, distinguitur.

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 14a-c) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, 31-62× $5-8 \mu \mathrm{~m}$; stipe extension aseptate, straight, 70-97 $\mu \mathrm{m}$ long, thick-walled with one basal septum, terminating in thin-walled, clavate to ellipsoidal vesicles (Fig. 14df), 6-8 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig. $14 \mathrm{~g}-\mathrm{i}$ ) with primary branches aseptate, $9-19 \times 3-5 \mu \mathrm{~m}$, secondary branches aseptate, $9-11 \times 4 \mu \mathrm{~m}$, each terminal branch producing $2-4$ phialides; phialides cymbiform to cylindrical, hyaline, aseptate, $10-15 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores (Fig. 14j-k) in moderate numbers, comprising of a septate stipe, and primary branches terminating in 2-3 phialides; primary branches straight, hyaline, $0-1$-septate, $16-$ $40 \times 4 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $17-28 \times 3-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 141) cylindrical, rounded at both ends, straight, 1 -septate, ( $11-$ ) $12-14(-15) \times 2-4 \mu \mathrm{~m}$ (av. $=$ $13 \times 3 \mu \mathrm{~m}$ ), frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth margins, white, buff yellow (19 d) to umber (13i) (reverse); chlamydospores extensive throughout medium, arranged in chains; reaching 75 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - Brazil, Sao Paulo, Aracruz nursery, from Eucalyptus cutting, 1992, A.C. Alfenas, holotype CBSH20607, culture ex-type CBS 210.94=PPRI 4450=UFV $125=$ IMI 361579; Madagascar, Isoamala-Beraketa, Mount


Fig. 13 Cylindrocladiella pseudocamelliae. a-e. Penicillate conidiophores. $\mathbf{f}-\mathbf{i}$. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{n}$. Terminal vesicles. $\mathbf{0}-\mathbf{p}$. Subverticillate conidiophores.
Q. Conidia. Scale bars: $\mathrm{A}=50 \mu \mathrm{~m}, \mathrm{~B}=50 \mu \mathrm{~m}$ (apply to $\mathbf{c}-\mathbf{e}$ ), $\mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{g}-\mathbf{q}$ )

Tolongo, substrate unknown, 7 Mar. 1994, collector unknown, culture CBS 115610=CPC 909=Fox 409.

Notes - Morphologically C. pseudohawaiiensis is difficult to distinguish from C. hawaiiensis, and therefore phylogenetic inference is required. It can be distinguished from $C$. infestans by its smaller conidium dimensions and terminal vesicle
shape. Cylindrocladiella pseudohawaiiensis can also be distinguished from other species in the $C$. infestans complex by different unique fixed nucleotides for three loci: BTUB positions 127 (A) and 384 (G); HIS3 positions 23 (C), 29 (C), 33 (A), 77 (G), 283 (indel), 285 (C), 288 (A), 314 (T), 349 (T) and 463 (T); TEF-1 $\alpha$


Fig. 14 Cylindrocladiella pseudohawaiiensis. a-c. Penicillate conidiophores. d-f. Terminal vesicles. g-i. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{k}$. Subverticillate conidiophores. L. Conidia. $\mathrm{A}=20 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{c}$ ), $\mathrm{D}=10 \mu \mathrm{~m}$ (apply to $\mathbf{e}-\mathbf{l}$ )
positions 153 (T), 244 (T), 288 (T), 289 (A), 290 (T), 337 (C), 465 (A), 471 (G), 478 (T) and 482 (G).

Cylindrocladiella pseudoparva L. Lombard \& Crous, sp. nov. - MycoBank MB561672, Fig. 15.

Etymology - Named after its morphological similarity to Cylindrocladiella parva.

Cylindrocladiellae parvae morphologice valde similis, sed ramis primariis conidiophorum majoribus distinguitur.

Teleomorph unknown. Conidiophores monomorphic, penicillate, mononematous and hyaline. Penicillate conidiophores (Fig. 15a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, 31-86×5-9 $\mu \mathrm{m}$; stipe extension aseptate, straight, 111-164 $\mu \mathrm{m}$ long, thick-walled with one basal septum, terminating in thin-walled, clavate to ellipsoidal to pyriform vesicles (Fig. 15f-h), 5-7 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig.15i-k) with primary branches aseptate, $16-32 \times 3-6 \mu \mathrm{~m}$, secondary branches aseptate, $8-18 \times 3-$
$5 \mu \mathrm{~m}$, each terminal branch producing 2-4 phialides; phialides doliiform to cymbiform, hyaline, aseptate, 10$17 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette absent. Subverticillate conidiophores not observed. Conidia (Fig. 151) cylindrical, rounded at both ends, straight, 1 -septate, $16-18(-20) \times 2-4 \mu \mathrm{~m}(\mathrm{av} .=17 \times$ $3 \mu \mathrm{~m}$ ), frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth margins, white with buff yellow (19 d) centre, umber (13i) (reverse); chlamydospores extensive throughout medium, arranged in chains; reaching 50 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - The Netherlands, Apeldoorn, Paleis Het Loo, from soil, Apr. 2010, P.W. Crous, holotype CBS-H20608, culture ex-type CBS $129560=$ CPC 18149; New Zealand, South Auckland, Karaka, Karaka road, from Vitis riparia, 16 Apr. 2007, K. Paice, culture CBS 122594; Switzerland, Mohlin Canton, Basel, from root of Quercus


Fig. 15 Cylindrocladiella pseudoparva. a-e. Penicillate conidiophores. $\mathbf{f}-\mathbf{h}$. Terminal vesicles. $\mathbf{i}-\mathbf{k}$. Conidiogenous apparatus with conidiophore branches and phialides. L. Conidia. Scale bars: $A=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{e}$ ), $\mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{g}-\mathbf{l}$ )
sp., 16 Mar. 1994, L. Petrini, culture CBS 113624=CPC 752.

Notes - Cylindrocladiella pseudoparva can be distinguished from C. parva and C. stellenboschensis by having larger primary, and smaller secondary branches. However, phylogenetic inference will be required to accurately identify it. Cylindrocladiella pseudoparva differs from other species in the C. parva complex by unique fixed nucleotides in two loci: BTUB position 199 (G) and 358 (A); HIS3 position 226 (T), 302 (A), 372 (T) and 436 (C).

Cylindrocladiella queenslandica L. Lombard \& Crous, sp. nov. - MycoBank MB561682, Fig. 16.

Etymology - Named after Queensland, the state in Australia from where it was collected.

Cylindrocladiellae infestantis morphologice valde similis, sed conidiis minoribus, (9-)10.5-13.5(-15) $\times 2-4 \mu \mathrm{~m}$, distinguitur.

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 16a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, $41-82 \times 6-9 \mu \mathrm{~m}$; stipe extension aseptate, straight, 117$180 \mu \mathrm{~m}$ long, thick-walled with one basal septum, terminating in thin-walled, cylindrical to lanceolate vesicles (Fig. 16j-m), $5-8 \mu \mathrm{~m}$ wide. Penicillate conidiogenous apparatus (Fig. 16fi) with primary branches aseptate, $13-23 \times 3-7 \mu \mathrm{~m}$, secondary branches aseptate, $9-12 \times 2-4 \mu \mathrm{~m}$, each terminal branch

conidiophores. P. Conidia. Scale bars: $\mathrm{A}=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{c}$ ), $\mathrm{D}=$ $20 \mu \mathrm{~m}$ (apply to $\mathbf{e}, \mathbf{n}$ ), $\mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{g}-\mathbf{m}, \mathbf{o}-\mathbf{p}$ )
producing 2-4 phialides; phialides reniform to doliiform to cymbiform, hyaline, aseptate, $7-15 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores (Fig. 16n-o) in moderate numbers, comprising of a septate stipe, and primary branches terminating in 2-3
phialides; primary branches straight, hyaline, $0-1$-septate, $22-$ $50 \times 3-4 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $17-41 \times 2-6 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 16p) cylindrical, rounded at both ends, straight, 1 -septate, (9-)10.5-13.5
$(-15) \times 2-4 \mu \mathrm{~m}(\mathrm{av} .=12 \times 3 \mu \mathrm{~m})$, frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth to undulate margins, white with straw (21 d) tint in patches, buff yellow (19 d) to umber (13i) (reverse); chlamydospores moderate throughout medium, arranged in chains; reaching 90 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimen examined - Australia, Queensland, from soil, 18 June 2009, P.W. Crous, holotype CBS-H20609, culture ex-type CBS 129574=CPC 17568; Australia, Queensland, from soil, 18 June 2009, P.W. Crous, culture CBS 129575= CPC 17569.

Notes - Cylindrocladiella queenslandica can be distinguished from other species in the C. infestans complex based on its smaller conidia, and unique fixed nucleotides for three loci: BTUB position 201 (T); HIS3 positions $110(\mathrm{G})$ and 310 (G); TEF-1 $\alpha$ positions 35 (A) and $455(\mathrm{~T})$.

Cylindrocladiella stellenboschensis L. Lombard \& Crous, sp. nov. - MycoBank MB561671, Fig. 17.

Etymology - Named after the town from which this species was first collected, Stellenbosch, South Africa.

Cylindrocladiellae parvae morphologice valde similis, sed conidiis majoribus, $(14-) 17-19(-21) \times 2-4 \mu \mathrm{~m}$, distinguitur.

Teleomorph unknown. Conidiophores monomorphic, penicillate, mononematous and hyaline. Penicillate


Fig. 17 Cylindrocladiella stellenboschensis a-d. Penicillate conidiophores. e-i. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{m}$. Terminal vesicles. N. Conidia. Scale bars: $A=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{d}$ ), $\mathrm{E}=10 \mu \mathrm{~m}$ (apply to $\mathbf{f}-\mathbf{n})$
conidiophores (Fig. 17a-c) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, 37$65 \times 6-9 \mu \mathrm{~m}$; stipe extension aseptate, straight, 109$169 \mu \mathrm{~m}$ long, thick-walled with one basal septum, terminating in thin-walled, clavate to naviculate vesicles (Fig. 17h-j), 5$7 \mu \mathrm{~m}$ wide. Penicillate conidiogenous apparatus (Fig. 17d-g) with primary branches aseptate, $13-28 \times 3-5 \mu \mathrm{~m}$, secondary branches aseptate, $10-16 \times 3-6 \mu \mathrm{~m}$, each terminal branch producing $2-4$ phialides; phialides doliiform to cymbiform, hyaline, aseptate, $12-21 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores not observed. Conidia (Fig. 17k-n) cylindrical, rounded at both ends, straight, 1 -septate, (14-)17-19(-21)× $2-4 \mu \mathrm{~m}(\mathrm{av} .=18 \times 3 \mu \mathrm{~m})$, frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth margins, white with straw ( 21 d ) tint in patches, umber (13i) (reverse); chlamydospores extensive throughout medium, arranged in chains; reaching 60 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - South Africa, Western Cape Province, Stellenbosch, Stellenbosch Botanical Gardens, from leaf litter, 31 Aug. 1992, P.W. Crous, holotype CBSH20610, culture ex-type CBS 110668=CPC 517; Canada, Toronto, Queens Park North, from leaf litter, 24 Apr. 2008, P.W. Crous, culture CPC 15200; Switzerland, Therwil Canton, Basel, from root of Quercus sp., 16 Mar. 1994, L. Petrini, culture CBS 116170=CPC 753.

Notes - This species can be distinguished from C. parva by its larger conidia and shape of the terminal vesicle. Furthermore, collarettes are also present on its phialides, whereas these are rare or absent for C. parva. Cylindrocladiella stellenboschensis differs from other lineages in the C. parva complex by unique fixed nucleotides in one locus: BTUB position 112 (A), 162 (G), 172 (A), 268 (C), 352 (T), 361 (C), 366 (G), 370 (T), 371 (G), 378 (A), 382 (A), 396 (A) and 495 (C).

Cylindrocladiella thailandica L. Lombard \& Crous, sp. nov. - MycoBank MB561680, Fig. 18.

Etymology - Named after the country where it was collected, Thailand.

Cylindrocladiellae infestantis morphologice similis, sed extensionibus stipitis longioribus, 123-183 $\mu \mathrm{m}$.

Teleomorph unknown. Conidiophores dimorphic, penicillate and subverticillate, mononematous and hyaline. Penicillate conidiophores (Fig. 18a-d) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, $49-80 \times 5-9 \mu \mathrm{~m}$; stipe extension aseptate, straight, 123$183 \mu \mathrm{~m}$ long, thick-walled with one basal septum, terminating in thin-walled, cylindrical to lanceolate vesicles (Fig. 18i-m), 5-7 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig. 18eh) with primary branches aseptate, $11-24 \times 4-8 \mu \mathrm{~m}$, secondary
branches aseptate, $7-14 \times 2-5 \mu \mathrm{~m}$, each terminal branch producing $2-4$ phialides; phialides reniform to doliiform to cymbiform, hyaline, aseptate, $8-13 \times 2-4 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Subverticillate conidiophores (Fig. 18n-p) in moderate numbers, comprising of a septate stipe, and primary branches terminating in $2-3$ phialides; primary branches straight, hyaline, $0-1$-septate, $40 \times 3 \mu \mathrm{~m}$; phialides cymbiform to cylindrical, hyaline, aseptate, $19-38 \times 2 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette. Conidia (Fig. 18q) cylindrical, rounded at both ends, straight, 1 -septate, (13-)14-16(-18) $\times 2-4 \mu \mathrm{~m}$ (av. $=$ $15 \times 3 \mu \mathrm{~m}$ ), frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth margins, white with buff yellow (19 d) centre, umber (13i) (reverse); chlamydospores extensive throughout medium, arranged in chains; reaching 70 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - Thailand, Chiang Mai, from soil, Oct. 2010, P.W. Crous, holotype CBS-H20611, culture ex-type CBS 129571=CPC 18835; Chiang Mai, from soil, Oct. 2010, P.W. Crous, CBS $129570=$ CPC 18834; Chiang Mai, from soil, Oct. 2010, P.W. Crous, CPC 18831.

Notes - Morphologically, C. thailandica is similar to C. infestans and C. longistipitata, with the exception that the stipe extensions are longer than those of $C$. infestans but shorter than those of $C$. longistipitata. Cylindrocladiella thailandica can also be distinguished from other species in the $C$. infestans complex by different unique fixed nucleotides for two loci: BTUB position 160 (G); HIS3 positions 27 (C), 30 (A), 60-63 (indel), 70 (A) and 117 (A).

Cylindrocladiella variabilis L. Lombard \& Crous, sp. nov. - MycoBank MB561673, Fig. 19.

Etymology - Named after its highly variable vesicle morphology.

Cylindrocladiellae lageniformis morphologice similis, sed vesiculis divergentibus.

Teleomorph unknown. Conidiophores monomorphic, penicillate, mononematous and hyaline. Penicillate conidiophores (Fig. 19a-e) comprising a stipe, a penicillate arrangement of fertile branches, a stipe extension and a terminal vesicle; stipe septate, hyaline, smooth, 41-91×5$9 \mu \mathrm{~m}$; stipe extension aseptate, straight, 67-106 $\mu \mathrm{m}$ long, thick-walled with one basal septum, terminating in thin-walled, clavate to fusoid to ovoid vesicles (Fig. 19j-m), 5-10 $\mu \mathrm{m}$ wide. Penicillate conidiogenous apparatus (Fig. 19f-i) with primary branches aseptate, 12-23×3$7 \mu \mathrm{~m}$, secondary branches aseptate, $9-14 \times 3-6 \mu \mathrm{~m}$, each terminal branch producing 2-4 phialides; phialides doliiform to cymbiform, hyaline, aseptate, $7-17 \times 2-6 \mu \mathrm{~m}$, apex with minute periclinal thickening and collarette absent. Subverticillate conidiophores not observed. Conidia


Fig. 18 Cylindrocladiella thailandica. a-d. Penicillate conidiophores. e-h. Conidiogenous apparatus with conidiophore branches and phialides. im. Terminal vesicles. $\mathbf{n}-\mathbf{p}$. Subverticillate conidiophores. Q. Conidia. $\mathrm{A}=50 \mu \mathrm{~m}$ (apply to $\mathbf{b}-\mathbf{d}$ ), $\mathrm{F}=10 \mu \mathrm{~m}$ (apply to $\mathbf{e}-\mathbf{q}$ )
(Fig. 19n) cylindrical, rounded at both ends, straight, 1septate, (9-) $11-13(-14) \times 2-3 \mu \mathrm{~m}(\mathrm{av} .=12 \times 3 \mu \mathrm{~m})$, frequently slightly flattened at the base, held in asymmetrical clusters by colourless slime.

Culture characteristics - Colonies raised (convex), cottony, with smooth margins, white, umber (13i)
(reverse); chlamydospores extensive throughout medium, arranged in chains; reaching 60 mm after 7 days on MEA at $24^{\circ} \mathrm{C}$ in the dark.

Specimens examined - Australia, Queensland, Daydream island, Whitsundays Island Resort, from soil, 2 Aug. 2009, P.W. Crous, holotype CBS-H20612, culture ex-type CBS


Fig. 19 Cylindrocladiella variabilis. a-e. Penicillate conidiophores. $\mathbf{f}-\mathbf{i}$. Conidiogenous apparatus with conidiophore branches and phialides. $\mathbf{j}-\mathbf{m}$. Terminal vesicles. N. Conidia. Scale bars: $A=20 \mu \mathrm{~m}$ (apply to $\mathbf{b}$ ), $C=50 \mu \mathrm{~m}$ (apply to $\mathbf{d}$ ), $E=10 \mu \mathrm{~m}$ (apply to $\mathbf{f}-\mathbf{n}$ )
$129561=$ CPC 17505, CPC 17504; Australia, Queensland, Lake Barrine, from soil, 18 June 2009, P.W. Crous, culture CBS 129562=CPC 17563.

Notes - Cylindrocladiella variabile can be distinguished from C. lageniformis by the high variability of its terminal vesicle shape. This species does not produce subverticillate conidiophores, whereas C. lageniformis produces them in moderate numbers (Crous 2002).

Cylindrocladiella pseudoinfestans L. Lombard \& Crous, nom. nov. - MycoBank MB561684

Basionym: Nectricladiella infestans Crous \& C.L. Schoch, Studies in Mycology 45: 55. 2000.

Etymology - Named after its morphological similarity to C. infestans.

Notes - Cylindrocladiella pseudoinfestans is introduced as a new name for $N$. infestans in the genus Cylindrocladiella. Nectricladiella infestans was incorrectly linked to its purported anamorph, C. infestans (Schoch et al. 2000), to which it is morphologically similar. Cylindrocladiella pseudoinfestans can be distinguished from other species in the C. infestans complex by different unique fixed nucleotides for three loci: BTUB position 395 (A); HIS3 positions 22 (T), 41 (G), 47 (A), 50 (A), 72 (T) and 272 (C); TEF-1 $\alpha$ positions 268 (A), 272 (G), 478 (A) and 480 (C).

## Discussion

In this study, several Cylindrocladiella isolates from numerous hosts and countries collected over the past two decades were shown to include a number of novel species. These species were recognised using phylogenetic inference and, where possible, supported by morphological features. The taxonomic status of several phylogenetic species identified in this study remains unresolved due to either representation by only a single isolate (e.g. CBS 116095) or culture sterility (e.g. clade containing CBS 115673). Naming these novel species in the anamorph genus Cylindrocladiella and not the teleomorph genus Nectricladiella follows the "strict priority" option as applied by Gräfenhan et al. (2011), which continued the approach of Lombard et al. (2009, 2010a-c), and Schroers et al. (2011) of naming fungi in the Hyprocreales with the oldest generic name, irrespective of its morph typification. Consequently, the novel species found in this study were named in the genus Cylindrocladiella (Boesewinkel 1982) rather than in the teleomorph genus Nectricladiella (Schoch et al. 2000).

Five species complexes could be identified in this study based on phylogenetic inference supported by morphological characterisation. Although previous authors (Victor et al. 1998, Schoch et al. 2000, Crous 2002, van Coller et al. 2005) acknowledged the presence of species complexes in the genus Cylindrocladiella, their sample sizes were small. In our study, a larger sample size, obtained from various culture collections, allowed a multi-gene analysis to more clearly identify species complexes in Cylindrocladiella.

The Cylindrocladiella camelliae species complex was shown to consist of several phylogenetic species, four of which were described as C. longiphialidica, C. natalensis, C. nederlandica and C. pseudocamelliae. Each of these four new species was distinguished from C. camelliae and each other by the morphology and dimensions of conidia, subverticillate conidiophores and stipe extensions. Geographical distribution of the various species in the C. camelliae complex reflected the cosmopolitan nature of this group of fungi. Cylindrocladiella nederlandica and C. natalensis were isolated from diseased plant material, and C. pseudocamelliae and C. longiphialidica were only isolated from soil, and their significance as plant pathogens still needs to be determined.

Cylindrocladiella cymbiformis is a newly described species closely related to both $C$. novaezelandiae, as well as novel lineages in the newly identified C. elegans species complex. Cylindrocladiella cymbiformis is not a cryptic species in the C. elegans complex as it can be distinguished from both C. novaezelandiae and C. elegans by its larger conidium dimensions and shorter stipe extensions. All isolates in this study representing the C. elegans complex originated from South Africa, whereas C. cymbiformis is
described here from soil samples collected in Australia. Cryptic species were not resolved in the C. elegans complex as the cultures were sterile.

Past studies have presented evidence of cryptic speciation within C. infestans (Victor et al. 1998, Schoch et al. 2000, Crous 2002, van Coller et al. 2005). In an attempt to resolve taxa in this complex, a large sample of C. infestans senso lato isolates was included in this study. Based on phylogenetic inference and morphological characterisation, a total of 12 cryptic species were identified. Of these, eight were described as novel taxa. All eight of these newly named species may be regarded as phylogenetic species, as morphological characters are limited to distinguish them from each other. These species are recognised using the genealogical concordance phylogenetic species recognition (GCPSR) criteria (Taylor et al. 2000) based on DNA sequence data for the five loci used in this study. As has been done for other fungal groups (O’Donnell et al. 2004, Grünig et al. 2008, Pavlic et al. 2009, Lombard et al. 2010b), these species are chiefly characterised by fixed single nucleotide polymorphisms (SNPs).

Schoch et al. (2000) described Nectricladiella infestans as the teleomorph state of $C$. infestans sensu lato from an isolate collected in Madagascar that produced perithecia in culture. With additional sequence data and isolates, van Coller et al. (2005) showed this isolate represented a cryptic species distinct from C. infestans senso stricto. This was further supported by the phylogenetic inference in this study, and based on GCPSR, Nectricladiella infestans has been provided with a new name, C. pseudoinfestans.

Cylindrocladiella clavata, C. lanceolata and C. variabilis are newly described here, closely related to C. lageniformis. They can be distinguished from each other and C. lageniformis based on the absence of subverticillate conidiophores, terminal vesicle morphology and conidium dimensions. All three of these species, with the exception of C. lanceolata, are presently only known from soil samples collected in Australia. Cylindrocladiella lanceolata was also isolated from a diseased Eucalyptus cutting in South Africa, adding another Cylindrocladiella species recorded from that country (Crous et al. 1993, Crous et al. 1994, Crous 2002, van Coller et al. 2005).

Phylogenetic inference applied in this study also identified a number of cryptic species within a large sample of $C$. parva sensu lato isolates. Only two of these cryptic species could be named here, as most isolates were sterile. Cylindrocladiella stellenboschensis and C. pseudoparva are difficult to distinguish from each other or from C. parva by morphology alone. These two species are recognised as phylogenetic species described according to the GCPSR criteria using fixed SNPs.

Isolates of C. peruviana used in this study also included cryptic species that could not be named. As with the $C$.
elegans complex, isolates representing these cryptic species were sterile and their taxonomy remains unresolved.

Traditionally, DNA sequence data for the ITS and BTUB gene regions were used to explore the phylogenetic relationship between Cylindrocladiella spp. (Victor et al. 1998, Schoch et al. 2000). Van Coller et al. (2005) introduced HIS3 sequence data for this group of fungi, increasing the gene regions that provide the most valuable information on the relationships among Cylindrocladiella spp. Data for these three gene regions have been available only for a small sample of Cylindrocladiella isolates. This present study has attempted to address this problem and also introduced partial TEF-1 $\alpha$ gene region sequences for all known Cylindrocladiella spp. Phylogenetic analysis of the individual gene regions showed that the TEF- $1 \alpha$ gene region provided the best resolution to distinguish between Cylindrocladiella spp., followed by BTUB and HIS3. As was found with Calonectria spp. (Lombard et al. 2010b), the ITS and LSU gene regions provided limited information to distinguish between Cylindrocladiella spp.

Identification of a large number of cryptic species within the genus Cylindrocladiella based on phylogenetic inference and morphological comparisons, highlights how little attention this group has received in the past. Although Cylindrocladiella spp. are generally not regarded as important plant pathogens, correct identification is essential for disease control and biosecurity implications. This study has revealed the importance of combining morphological and phylogenetic data to understand the taxonomic issues surrounding this group of fungi.

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